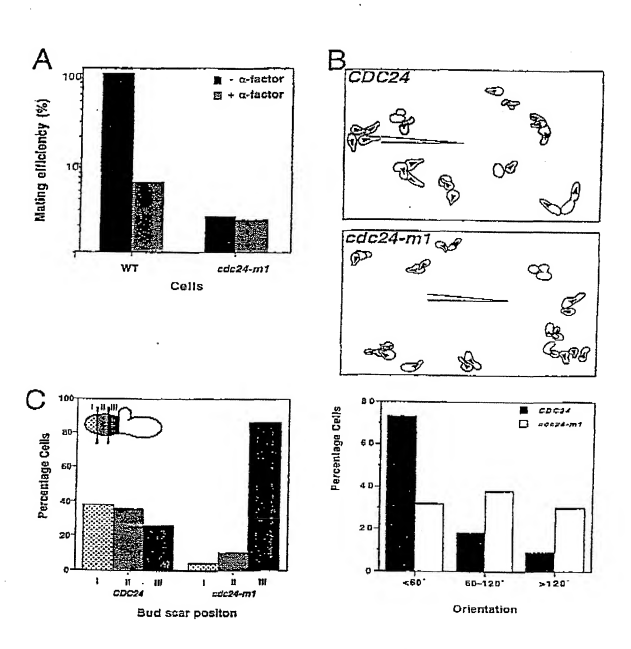
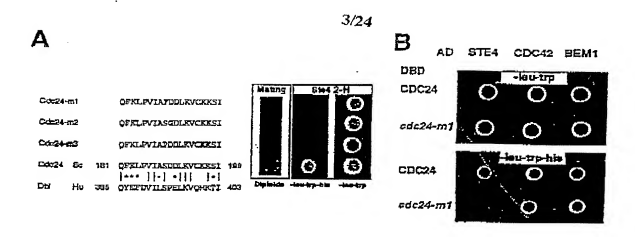


FIG. 1









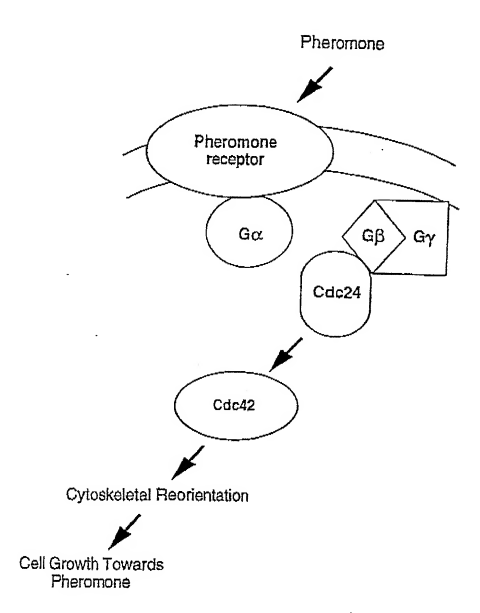
C

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Extracts Eluates

FIG. 3









# Figure 5

5																		
J	2113/1 ATG GAA CA								TTT		ACC							
	M E H 2173/21	P	P	A	A	L	R	T	도 220:	S 3/31	T	Q	S	T	5	Ş	L	N
10	TCA GTA AG S V S	T ACI	GTT V	TCG S	TĊT S	TCA S	AĠA R	ATT I	GTT V	TCT S	S	GGC G	CCA P	GTC V	AAT N	ATA I	aac N	AAT N
	2233/41 TTC AAT AA	A CCA	AGT	ACT	ccc	λλλ	GAC	CAT	226. TTA	3/51 TTC	<b>ተ</b> ልተ	CGA	ጥርም	CAA	ጥሮል	ሮሞአ	አክክ	<u>ሶር</u> አ
	F N K	₽	5	T	₽	к	D	H	L	F	Y	R	c	E	S	L	K	R
15	2293/61 AAA CTA CA K L Q	A AAA K	ATC	CCT	GGC G	ATG M	GAA E	CCA P	TTT	3/71 TTG L	AAC N	CAA Q	GCT A	TTC F	AAT N	CAG Q	GCT A	GAA E
	2353/81		<i>-</i> 23.3	<i>-</i>				-	238:	3/91								_
20	CAA CTC AG Q L S 2413/101	T GAA	Q Q	Q	GCA A	L L	GCT A	TTG L	A	CAG Q 3/11:	E	AGA R	AGC S	AAT N	GGA G	AAT N	GGA G	CAT H
	AGT AAT GG S N G	C AAA K	CGT R	CAT H	CAA Q	TCA S	TTA L	GAC D	ĠĠT G	GCC A	ATG M	AAT N	AGA R	CTT L	TCA \$	GTT V	GĞT G	TCT S
25	2473/121 GAT AGT AG	ד דרפ	ATĊ	ĊΑΑ	ርጎርትንግ	ΤĊΑ	тта	ACA	250:	3/13: ATC	GCG.	ACC	ייבב	aca	ጥሮክ	א כיכי	TON	rp/Drp
	D S S 2533/141	Ş	I	Q	G	S	L	T	R 256:	M 3/15:	A 1	Т	И	A	s	T	s	s
20	TTA ATC AG	r GGT G	ATG M	CCA P	AAC N	aac N	AAC N	ACT T	Ľ	F	T	TTT F	ACT T	GCA A	GGG G	GTT V	TTA L	CCA P
30	2593/161 GCT AAT AT A N I	r agt s	GTC V	GAT D	ÇÇT P	GCT A	AÇÇ T	ÇAT H	CTT	3/17: TGG W	AAA	TTG L	TTC F	CAA Q	CAA O	GGG G	GCÇ A	ÇCÇ
	2653/181 TTT TGT GT	-							2683	3/19:	1		_		_			-
35	F C V	L	ĭ	Ŋ	H	Ï	Ļ	Þ	D	s	Õ	I	P	V	V	AGT S	TCT \$	D D
	2713/201 GAC TTG AG D L R	A ATT	TGC C	AAA K	AAA K	TCA S	GTA V	TAT Y	GAC D	3/21: TTT F	TTA L	ATT I	GCC A	GTC V	AAG K	ACA T	CAA	TTG L
40	2773/221 AAT TTT GA								TCC		GTT						CAA	GAT
	N F D 2833/241	D	E	N	М	F	T	I	S 2863	/25	V L	F	S	D	N	A	Q	D
45	TTA ATC AAC L I K 2893/261	I ATT	ATT	GAT D	GTC V	ATT	AAT N	AAA K	L		A.		TAC Y	TCA S	gat D	GCT A	agt S	GAC D
70	CTG GGT GGT	GGC G	GAT D			GTA V	TAA N	ATG M	GAT	A GŁŁ	CAA Q	TTA	ACC T	GAT D	gaa E	AGA R	TCA S	AAA K
	2953/281 GTT TTC CG/	GAA	ATT	ATC	ĠAA	AĊA	GAA	AGA	2983 AAA	3/293 TAT	CTT	CAA	GAC	TTC	GAA	cπΔ	ATC	TET
50	V F R 3013/301	E	I	I	E	r	E	R	ĸ		v		D	L	E	L	М	Ç
	AAA TAC CGT	CAA Q	GAT D	CTA L	ATT I	GAA E	GCC A	GAA E	AAT N	TTG	TCT S	TCA S	GAA E	CAA Q	ATT I	CAC H	TTG L	TTA L
55	3073/321 TTC CCA AAT	TŤA	AAT	GAG	ATT	ATT	GAT	TTT	3103 CAA	1/331 AGA	L CĠA	TTC	כידיכי	ΔAT'	GGG	ጥጥል	CA A	יייבאיי
	F P N 3133/341	L	N	E	I	1	Ď	F	Q	R (/35)	R	F	L	N	G	Ļ	Ē	Ĉ
	AAC ATC AAT	' GTA V	CCT P	ATT I	AGA R		CAA Q	AGA R	ATT	GGA	TCA	GTA V		ATT I		GCT A	TCT S	TTG L
60	3193/361 GGC CCT TTC								3223	1/373	L		_					
	G P P 3253/381	N	A	Y		D	M	Ţ	I 3283	G	Q	L		A		D	TTG L	I
65	AAC AAA GAA N K É	GCT A	GCT A	AAT N	TTG L	AAA K	AAA K	TCG S	TCA	AGT S	CTA	CTT L				TTT F	GAA E	CTT L
	3313/401 CAA TCG TAT	ATA	TTA	AAG	cce	ATC	CAA		3343 TTG	/413 TGT	L AAA	TAC	CCA	CTT	TTG	TTG	AAA	GAG
	Q S Y 3373/421	-		K		I		R	L 3403	/431	Į						K	Ê
70	TTA ATC AAA L I K	ACA T	TCA S	CCA P	GAA E	TAT Y	TCA S	AAA K	CAG Q				GĠĊ G	AGC S	TCG S	TCA S	TCG S	ACA T
	3433/441	_							3463	/451	L	•	-		-	_		_
	TCA TTC AAT	GAA E	TTA L	TTG L	V GIG	GCT A	AAA K	ACT T	GCA A	ATG M	AAA K	GAA E	TTG L	GCA A	AAT N	CAA	GTC V	AAT N



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	3493/461								352	3/47	1							
	GAG GCG CAA	AGA	. CGA	. GCA	GAA	AAT	ATC	GAA	CAT	TTG	GAA	. AAA	. CTA	. AAA	GAA	AGA	GTA	GGT
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_	3553/481								358	3/49	1							
5	AAT TGG CGT	GGG	TTT	AAT	TTG	GAT	GCT	CAA	GGA	GAA	CTA	TTA	TTC	CAC	GGA	CAA	GTT	GGG
	N W R	G	F	N	L	Þ	A	Q	Ġ	E	Ŀ	Ľ	F	H	G	Q	v	G
	3613/501								364	3/51	l							
	GTT AAA GAT	GCT	GAA	AAT	GAA	AAG	GAA	TAC	GTT	GCT	TAT	CTT	TTT	GAA	AAA	ATC	GTA	TTT
	V K D	A	E	N	E	K	E	Y	v	A	Y	L	F	E	K	I	v	F
10	3673/521								370	3/53	1							*
	TTT TTC ACA	. GAA	ATT	GAT	GAT	ACC	AAA	AAA	TCT	GAT	AAA	CAG	GAA	AAG	AAG	AGC	AAG	TTT
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	3733/541								376	3/55	1							_
	TCG ACA AGA	AAG	AGA	TCA	ACT	TCA	TCA	AAT	CTT	AĞT	TCA	TĊG	ACT	ACT	AAT	TTG	TTG	GAA
15	S T R	K	R	\$	T .	5	S	N	T.	5	5	5	T	T	N	L	L	E
	3793/561								3823	3/57	1						_	_
	TCA ATA AAC	AAT	TCC	ÇÇA	AAG	GAT	AAC	ACA	TTG	CÇA	TTG	GAA	TTA	AAG	GGA	AGA	GTT	TAT
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~~	3853/581								3883	3/59	7							
20	ATA TCG GAG	ATT	TAT	AAC	ATT	TCÇ	ĢÇA	CCA	AAC	ACT	CCT	GGC	TCA	ACT	CTA	ATC	ATC	TCA
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	TGG TCA GGT W S G	R	K	E	3	G	S S	F	ACT			TAT						
25	3973/621	IX.	K	-	-7	-	۵				R	X	R	\$	£	E	A	R
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	4033/641	_		•	_		_	~		, 765:		<i>I</i>	1-1	14	~	Q	I	H
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30	к к ь	R	D	S	D	S	S	F	N		D	D	5	A	I	Y	D	Y
	4093/661							-,	4123	/67	1							_
	ACG GGT ATT	AGT	ACG	TCA	CCA	GTC	AAT	CAA	TCA	ACT	CAA	CAA	CAA	TAC	TAT	CAT	CAT	CCC
	T G I	S	${f T}$	S	₽	v	N	Q	ន		Q	Q	Q	Y	Y	D	H	R
	4153/681							-	4183	/693	1	•						
35	GGC TCT CAC	AGT	TCC	ÇÇC	CAT	CAC	TCA	TCG	TCA	TCC	ACT	TTG	AGT	ATG	ATG	AAG	AAT	AAT
	G S H	S	S	R	H	H	S	S	ន	S	${f T}$	L	s	M	M	K	N	N
	4213/701								4243	/713	L							
	AGA GTT AAA	T.C.T.	GGT.	GAT	TTG	AGT	AGA	ATA	TCT	TCA	ACT	TCA	ACA	ACA	TTA	GAT		
40	R V K 4273/721	S	G	D	Ļ	S	R	I	S			S	T	T	L	D	S	F·
70	AGT AAC AAC	בותים	יתעע	ccc	ጥሮክ	CCN	<b>አአ</b> ም	700	4303	7731		mam			-			
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	43 <i>9</i> 3/761								4423	1777						_	_	_
	TCA GAG CCA	TTG	ATT	GTC	TAA	GÇA	ÇAA	ATT	GAG	TAT	AAT	GAC	CTT	ATT	CAG	AAA	ATT	ATC
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	4453/781								4483	/791								• •
50	TCC CAG ATT	ATC .	ACT	TCG .	AAC	TTG	GTĢ	GCA	GAT	GAT	GTÇ	AAT	ATT	AGT	CGA	TTG .	AGA	TAT
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	4513/801	~~~	caca y	<b>030</b>	mmm	ama	7 7 m	mmer	4543	/811								
	AAA GAC GAC	GAA :	GGA G	GAC D	F	V V	N NAT	TTG	AAT	TCA	GAT							
55	4573/821	2	<b>.</b>		F	v	W	L		\$ /071		D	D	W	G :	Ľ	V	L
22	GAT ATG TTA	ACC	AGT :	GAA	CAC	יויידידי	<b>ጉ</b> ልሮ	C22	4603	ውርል Leat	7/7/7	7. 7. CT	~~~	777	~~»			
	D M L	T	s :	E :	D	F	Y	Q	T :	s .								
	4633/841	_ '		•		-	-	~	-		J	7.4	متد	IC.	R.	s ·	V	T
	GTG TGG GTT	TCT	TGA															
60	V W V S *																	



# Figure 6

		•	
5	Blastp line up	of S.c. Cdc24p and C.a. Cdc24p	
	8.c. Cdc24p; 1	MAIQTR-FA M R F+	8
10	C.a. Cdc24p: 1	MEHPPAALRIFSTO	14
	5.c. Cdc24p: 9	SGTSLSDLKPKPSATSISIPMONVMNKPVTEQDSLFHICANIRKRLEVLPOLKPFLQI	L 66
	C.a. Cdc24p: 15	STSSLNSVSTVSSSRIVSSGPVNINNFNKPSTPKDHLFYRCESLKRKLQKIPGMEPFLNC	74
15	S.c. Cdc24p: 67	AYQSSEVLSERQSLLLSQKQHQELLKSNGANRDSSDLAPTLRSSSISTATSLMSMEC	3 123
	C.a. Cdc24p: 75	APNQAEQLSEQQALALAQERSNGNGHSNGKRHQSLDGAMNRLSVGSDSSSIQGSLTRMAT	r 134
20	S.c. Cdc24p: 12	4 ISYTNSNPSATPNMEDTLLTFSMGILPITMDCDFVTQLSQLFQQGAPLCILFNSVKPQFK + T+S S PN +TL TF+ G+LP + DP T L +LFQQGAP C+L N + P +	183
	C.a. Cdc24p: 13	THE TOUBLE OF THE THE THE TAG V DEAN IS V DEATHLWKL F QQ QAPF CV LINHILPDS Q	193
	S.c. Cdc24p: 18	4 LPVIASDDLKVCKKSIYDFILGCKKHPAFNDEELFTISDVFANSTSQLVKVLEVVETLMN +PV++SDDL++CKKS+YDF++ V	7 243
25	C.a. Cdc24p: 19	4 IPVVSSDDLRICKKSVYDFLIAVKTQLNFDDENMFTISNVFSDNAQDLIKIIDVINKLLA	253
	S.C. Cdc24p: 24:	4 SSPTIFDCVCVCOATAGE TOTAGE	
30	C.a. Cdc24p: 25	4 EYSDASDSGGGDEDVNMDVQITDERSKVFREIIETERKYV DLE++ KY	302
	s.c. Cdc24p: 304	RQQLLDSNLITSEELYMLFPNLGDAIDFQRRFLISLEINALVEPSKQRIGALFMH-SKHF RQ L+++ ++SE++++LFPNL + IDFQRRFL LE N V QRIG++F+H S	362
2.5	C.a. Cdc24p: 303		362
35	S.c. Cdc24p: 363	FKLYEPWSIGQNAAIEFLSSTLHKMRVDESQRFIINNKLELQSFLYKPVQRLCRYPLLVK F YEPW+IGQ AI+ ++ ++ ++ ELQS++ KP+QRLC+YPLL+K	422
	C.a. Cdc24p:.363	FINALEPWITGQLTAIDLINKEAANLKKSSSLLDPGFELQSYILKPIQRLCKYFLLLK	419
40	C.a. Cdc24p: 423	ELLAESSDDNNTKELEAALDISKNIARSINENQRRTENHQVVKKLYGRV EL+ SS + EL A K +A +NE QRR EN + ++KL RV ELIKTSPEYSKQDPHGSSSSTSFNELLVAKTAMKELANQVNEAQRRAENIEHLEKLKERV	471
	S.c. Cdc24p: 472	VNWKGYRISKFGELLYFDKVFISTTNSSSEPEREFEVYLFEKII1LFSEVVTKKSASSLI	479
45	C.a. Cdc24p: 480	NW+G+ + GELL+ +V + +E E+E+ YLFEKIIILFSEVVTKKSASSLI GNWRGFNLDAQGELLFHGQVGVKDAENEKEYVAYLFEKIVFFFTEIDDTKKSDKQE	531
	S.c. Cdc24p: 532	LKKKSSTSASISASNITDNNGSPHHSYHKRHSNSSSSNNIULGGGGARAITHGGTNOO	535
. 50	C.a. Cdc24p: 536	K K ST ++\$N+  KKSKFSTRKRST\$\$NL	591
. 50	S.c. Cdc24p: 592	NSMNSSSSLFKL\$ANEPKLDLRGRIMIMNINOTIDON NDSINIBULGI KRO	540
	C.a. Cdc24p: 569	N+ L+L+GR+ I + I N +L I+W KE G+F L+ NTLFLELKGRVYISEIYNISAPNTPGSTLIISWSGRKESGSFTLR	613
55	S.c. Cdc24p: 650	FKNEETRDNWSSCLQQLIHDLKNEQFKARHHSSTSTTSSTAKSSSMMSPTTT	701
	C.a. Cdc24p: 614	YRSEEARNQWEKCLRDLKTNEMNKQIHKKLRDSDSSFNTDDSAIYDYTGISTSPVNQSTO	673
60	S.c. Cdc24p: 702	MNTPNHHNSRQTHDSMASFSSSHMKRVSDVLPKRRTTSSSFESEIKS	748
	C.a. Cdc24p: 674	QQYYDHRGSHSSRHHSSSSTLSMMKNNRVKSGDLSRISSTSTTLDSFSNNLNGSPNTTNP	733
	S.c. Cdc24p: 749	ISENPKNSIPESSILFRISYNNNSNNTSSSEIFTLLVEKVWNFDDLIMAINSKISN	804
65	C.a. Cdc24p: 734	SLMSSDATKTIPTFDVAIKLLYKSTELSEPLIVNAQIEYNDLLQKIISQIITSN	787
	S.c. Cdc24p: 805	THNNNISPITKIKYODEDGDEVVIGGDEDMINIA KEMI A ENDYEMINIA	
	C.a. Cdc24p: 788	++++ I++++Y+D++GDFV L SD+DW + +ML + F + LVADDVN-ISRLRYKDDEGDFVNLNSDDDWGLVLDMLTSEDFYQTSSNEKRSVTVWVS	344
70		- #2100111111011VWV3 (	/ <del>* *</del>



Figure 7a

8/24

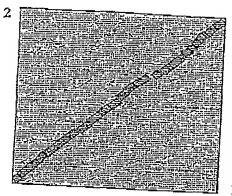
# BLAST 2 SEQUENCES RESULTS VERSION BLASTP 2.0.9 [May-07-1999]

Matrix 0.BLOSUM62 gap open: 11 gap extension: 1 x\_dropoff: 50. expect: 10.0 wordsize: 3 Filter

Sequence 1 Icl|S.c. Cdc24p Length 854 (1..854)

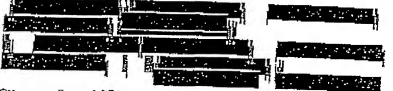
Sequence 2 lcl/C.a. Cdc24p Length 844 (1..844)





NOTE: The statistics (bitscore and expect value) is calculated based on the size of nr database

Score = 446 bits (1136), Expect = e-124 Tdentities = 288/881 (32%), Positives = 464/881 (51%), Gaps = 112/881 (12%)





SGTSLSDLKPKPSATSISIPMQNV--MNKPVTEQDSLFHICANIRKRLEVLPQLKPFLQL 66 Query: 9 NKP T +D LF+ C +++++L+ +P ++PFL <sup>‡</sup>3bjcz: 15

STSSLNSVSTVSSSRIVSSGPVNINNFNKPSTPKDHLFYRCESLKRKLQKIPGMEPFLNQ 74

Query: 67 AYQSSEVLSERQSLLLSQKQHQELLKSNGANRDSSDLAP---TLRSSSISTATSLMSMEG 123

sbjct: 75 AFNQAEQLSEQOALALAQERSNGNGHSNGKRHQSLDGAMNRLSVGSDSSSIQGSLTRMAT 134

Query: 124 ISYTNENPSATENMEDTLUTFSMGILPITMDCDPVTQLSQLFQQGAPLCILFNSVKPQFK 183 + T+S S PN +TL TF+ G+LP + DP T L +LFQQGAP C+L N + P

Sbjct: 135 nastsslisgmpn-nntlftftagvlpanisvdpathlwklfqqgapfcvlinhilpdsq 193

Query: 184 LPVIASDDLKVCKKSTYDFILGCKKHFAFNDEELFTISDVFANSTSQLVKVLEVVETLMN 243 F+DE +FTIS+VF+++

Sbjct: 194 IPVVSSDDLRICKKSVYDFLIAVKTQLNFDDENMFTISNVFSDNAQDLIKIIDVINKLLA 253

Query: 244 SSPTIFPSKSKTQQIMNAENQERHQPQQSSKKHNEYVKIIKEFVATERKYVHDLEILDKY 303

+ +E K+ +E + TERKYV DLE++ KY Sbjec: 254 EYSDASDSGGGDEDV------NMDVQITDERSKVFREIIETERKYVQDLELMCKY 302

Query: 304 RQQLLDSNLITSEELYMLFPNLGDAIDFQRRFLISLEINALVEPSKQRIGALFMH-SKEF 362 RO L+++ ++SE++++LFPNL + IDFQRRFL LE N V

Sbjct: 303 RQDLTEAENLSSEQIHLLFPNLNEIIDFQRRFLNGLECNINVPIRYQRIGSVFIHASLGP 362

Query: 363 FKLYEPWSIGQNAAIEFLSSTLHKMRVDESQRFIINNKLELQSFLYKPVQRLCRYPLLVK 422 +++

Sbjct: 363 FNAYEPWTIGQLTAIDLINKEAANLKKSSS-ELQS++ KP+QRLC+YPLL+K --LLDPGFFT.nevi



# F Figure 7a cont.

```
Query: 423 ELLAE-----SSDDNNTKELEAALDISKNIARSINENQRRTENHQVVKKLYGRV 471
                       SS + EL A K +A +NE QRR EN + ++KL RV
     SDJCE: 420 ELIKTSPEYSKODPHGSSSSTSFNELLVAKTAMKELANQVNEAQRRAENIEHLEKLKERV 479
     Query: 472 VNWKGYRISKFGELLYFDKVFISTTNSSSEPEREFEVYLFEKIIILFSEVVTKKSASSLI 531
                NW+G+ + GELL+ +V + +E E+E+ YLFEKI+ F+E+ K +
     Sbjct: 480 GNWRGFNLDAQGELLFHGQVGV----KDAENEKEYVAYLFEKIVFFFTEIDDTKKSDKQE 535
     Query: 532 LKKKSSTSASISASNITDNNGSPHHSYHKRHSNSSSSNNIHLSSSSAAAIIHSSTNSSDN 591
    Query: 592 NSNNSSSSSLFKLSANEPKLDLRGRIMIMNLNQIIPQN--NRSLNITWESIKEQGNFLLK 649
                               L+L+GR+ I + I N +L I+W KE G+F L+
    Sbjc: 569 NT-----LPLELKGRVYISEIYNISAPNTPGSTLIISWSCRKESGSPTLR 613
    Query: 650 FKNEETRDNWSSCLOQLIHDLKNEQFKARHHSSTSTTSS----TAKSSSMMSPTTT 701
    Sbjet: 614 YRSEEARNQWEKCLRDLKTNEMNKQIHKKLRDSDSSFNTDDSAIYDYTGISTSPVNQSTQ 673
   Query: 702 MNTPNHHNSRQT--HDSMASFSSSHMKRVS----DVLPKRRTTSSSFESEIKS----- 748
   Sbjct: 674 QQYYDHRGSHSSRHHSSSSTLSMMKNNRVKSGDLSRISSTSTTLDSFSNNLNGSPNTTNP 733
   Query: 749 -- ISENFKNSIPESSILFRISYNNNSNNTSSSEIFTLLVEKVWNFDDLIMAINSKI--SN 804
               +$ + +IP + ++ Y +T SE L+V
   Sbjct: 734 SLMSSDATKTIPTFDVAIKLLY---KSTELSE--PLIVNAQIEYNDLLQKIISQIITSN 787
   Query: 805 THNNNISPITKIKYQDEDGDFVVLGSDEDWNVAKEMLAENN 845
                ++++ I++++Y+D++GDFV L SD+DW + +ML
   SDJC: 788 LVADDVN-ISRLRYKDDEGDFVNLNSDDDWGLVLDMLTSED 827
  CPU time: 0.26 user secs.
                                    0.02 sys. secs 0.28 total secs.
  Gapped
  Lambda
     0.270 0.0470 0.230
  Matrix: BLOSUM62
  Gap Penalties: Existence: 11, Extension: 1
  Number of Hirs to DB: 12253
  Number of Sequences: 0
  umber of extensions: 709
Number of successful extensions: 15
 Number of sequences better than 10.0: 1
 Number of HSP's better than 10.0 without gapping: 1
 Number of HSP's successfully gapped in prelim test: 0
 Number of HSP's that attempted gapping in prelim test: 0
 Number of HSP's gapped (non-prelim): 1
 length of query: 844
 length of database: 90,077,593
 effective HSP length: 63
 effective length of query: 781
 effective length of database: 83353792
 effective search space: 65099311552
 effective search space used: 65099311552
 ጉ: 9
A: 40
X1: 16 ( 7.2 bits)
X2: 128 (49.9 bits)
X3: 128 (49.9 bits)
S1: '42 (21.9 bits)
$2: 73 (32.8 bits)
```



Figure 7b

10/24

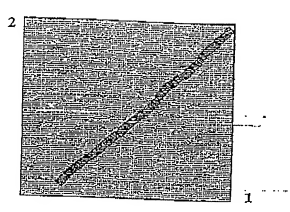
# BLAST 2 SEQUENCES RESULTS VERSION BLASTP 2.0.9 [May-07-1999]

Matrix OBLOSUM62 gap open: 11 gap extension: 1 x\_dropoff: 50 expect: 10.0 wordsize: 3 Filter Align

Sequence 1 Icl|S.c. Cdc24p Length 854 (1..854)

Sequence 2 Icl|S.p. Cdc24p Length 834 (1..834)





NOTE: The statistics (bitscore and expect value) is calculated based on the size of nr database

Score = 238 bits (601), Expect = 1e-61 Identities = 212/760 (27%), Positives = 348/760 (44%), Gaps = 94/760 (12%)



Query: 156 DPVTQLSQLFQQGAPLCILFNSVKPQPKLPVIASDDLK---VCKKSIYDFILGCKKHFAF 212 + G PLC LFN + + KL V +S L+ VCK S+Y F+L CK DPVT+÷

Sbjet: 67 DPVTEIWLFCRLGYPLCALFNCLPVKQKLEVNSSVSLENTNVCKASLYRFMLMCKNELGL 126

Query: 213 NDEELFTISDVFANSTSQLVKVLEVVETLMNSSPTIFPSKSKTQQIMNAENQERHQPQQS 272

D LF+IS+++ ST+ LV+ L+ +2 L+

SDJct: 127 TDAALFSISEIYKPSTAPLVRALQTIELLLKKYEVSNTTKSSSTPSPSTDDNVPTGTLNS 186

Query: 273 SKKHNEYVKIIKEFVATERKYVHDLEILDKYRQQLLDSNLITSPELYMLFPNLGDAIDFQ 332 ++ E TE KY+ DLE L Y L ++++

Sbjcc: 187 LIASGR--RVTAELYETELKYIQDLEYLSNYMVILQQKQILSQDTILSIFTNLNEILDFQ 244

Query: 333 RRFLISLEINALVEPSKQRIGALFMHSKHFFKLYEPWSIG-QNAAIEFLSSTLHKMRVDE 391 RRFL+ LE+N + +QR+GALF+ + F +Y+ +

5bjct: 245 RRFLVGLEMNLSLPVEEQRLGALFIALEEGFSVYQVFCTNFPNAQQLIIDNQNQLLKVAN 304

Query: 392 SQRFIINNKLELQSFLYKPVQRLCRYPLLVKELL-AESSDDNNTKELFAALDISKNIARS 450

EL + L KP+QR+C+YPLL+ +LL S

Sbjct: 305 ----LLEPSYELPALLIKPIQRICKYPLLLNQLLKGTPSGYQYEEELKQGMACVVRVANQ 360

Query: 451 Inengratenhovvkklygrvvnwkgyriskfgellyfdkvpisttnssseperefevyl 510

+NE +R EN + +L RV++WKGY + FG+LL +D V + ++ ERE+ VYL Sbjct: 361 VNETRRIHENRNAIIELEQRVIDWKGYSLQYFGQLLVWDVVNV----CKADIEREYHVYL 416

Query: 511 FEKIIILFSEVVT-KKSASSLILKKKSSTSAS-----ISASNITDN------ 550 FEKI++ E+ T K+ A S+ + KK+

I SNIT Sbjet: 417 FEKILLCCKEMSTLKROARSISMNKKTKRLDSLOLKGRIT



### | Figure 7b cont.

```
Query: 551 --NGSPHHSYHKRHSNSSSSNNIHLS------SSSAAAIIHSSTNSSDNNSNNSSSS 599
G P H + S+ ++S + I S+ ++ N N SSS
    Sbjct: 477 FWRGDPQHESFILKLRNEESHKLWMSVLNRLLWKNEHGSFKDIRSAASTPANPVYNRSSS 536
    Query: 600 SLFKLSANEPKLD-LRGRIMIMNLN---QIIPQNNRSLNITWESIKEQGNFLLKFKNEET 655
                 K N D LR + N+N I +++S T + K+
    Sbjct: 537 QTSK-GYNSSDYDLLRTHSLDENVNSPTSISSPSSKSSPFTKTTSKDT-----KSATT 588
    Query: 656 RDNWSSCLQQLIHDLKNEQFKARHHSSTST----TSSTAKSSSMMSPTTTMNT--PNHH 708
                 S +L + R +TST +SSTA S +S + +N+ +++
   sbjct: 589 TDERPSDFIRLNSEESVGTSSLRTSQTTSTIVSNDSSSTASIPSQISRISQVNSLLNDYN 648
   Query: 709 NSRQTH------DSMASF---SSSHMKRVSD-------VLPKRRTTSSSFESE 745
                              S++ F SSS +++ D + P++ + S+ +5+
   Sbjct: 649 YNRQSHITRVYSGTDDGSSVSIFEDTSSSTKQKIFDQPTTNDCDVMRPRQYSYSAGMKSD 708
   Query: 746 IKSISENFKNSIPESSILFRISYNNNSNNTSSSEI----FTLLVEKVWNFDDLIMAINSK 801
                      S+ SS +S N +N + L+V FD+L+ + K
  Sbjct: 709 GSLLPSTKHTSLSSSSTSTSLSVRNTTNVKIRLRLHEVSLVLVVAHDITFDELLAKVEHK 768
  Query: 802 IS--NTHNNNISPITKIKYQDEDGDFVVLGSDEDWNVAKE 839
                     + ++KY DEDGDF+ + SDED +A E
   ojec: 769 IKLCGILKQAVPFRVRLKYVDEDGDFITITSDEDVLMAFE 808
  CPU time: 0.26 user secs.
                                   0.04 sys. secs
                                                          0.30 total secs.
  Gapped
  Lambda
            0.0470
                     0.230
 Matrix: BLOSUM62
 Gap Penalties: Existence: 11, Extension: 1
 Number of Hits to DB: 10384
 Number of Sequences: 0
 Number of extensions: 671
 Number of successful extensions: 13
 Number of sequences better than 10.0: 1
 Number of HSP's better than 10.0 without gapping: 1
Number of HSP's successfully gapped in prelim test: 0
Number of HSP's that attempted gapping in prelim test: 0
Wimber of HSP's gapped (non-prelim): 2
 "juth of query: 834
length of database: 90,077,593
effective HSP length: 61
effective length of query: 773
effective length of database: 83489227
effective search space: 64537172471
effective search space used: 64537172471
r: 9
A: 40
15 ( 7.3 bits)
(2: 128 (49.9 bits)
(3: 128 (49-9 bits)
31: 41 (21.7 bits)
32: 73 (32.8 bits)
```



Figure 8

- 5 Sc KLPVIASDDLKVCKKSIYDFIL (SEQ ID No 25) ++PV++SDDL++CKKS+YDF++ Ca QIPVVSSDDLRICKKSVYDFLI (SEQ ID No 26)
- Sc = Saccharomyces cerevisiaeCa = Candida albicans



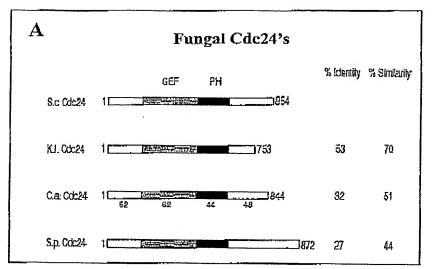
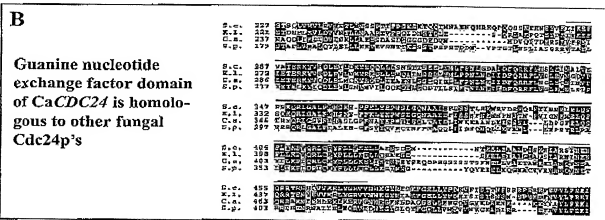
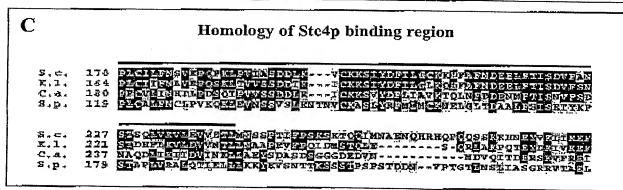
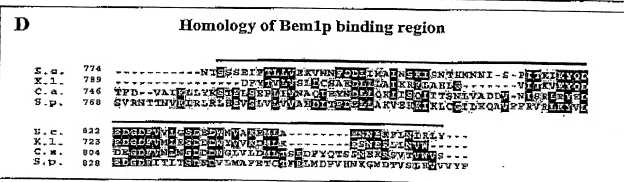
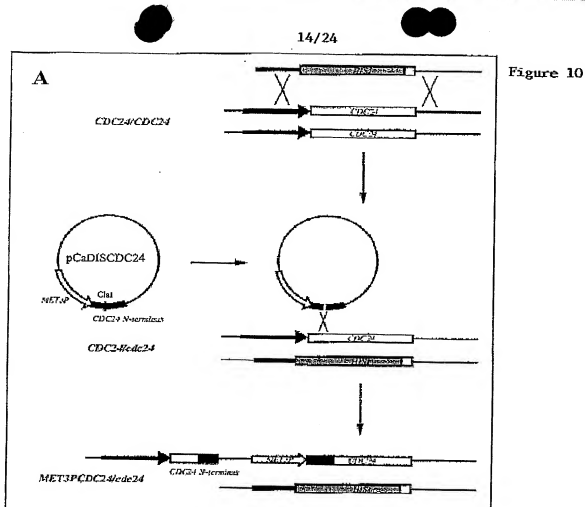


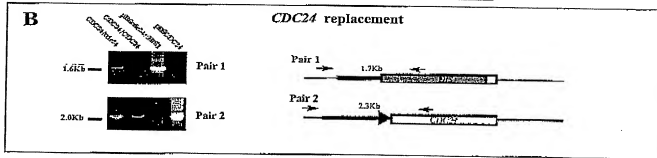
figure 9











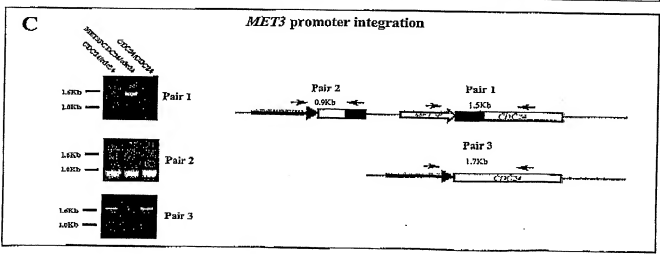
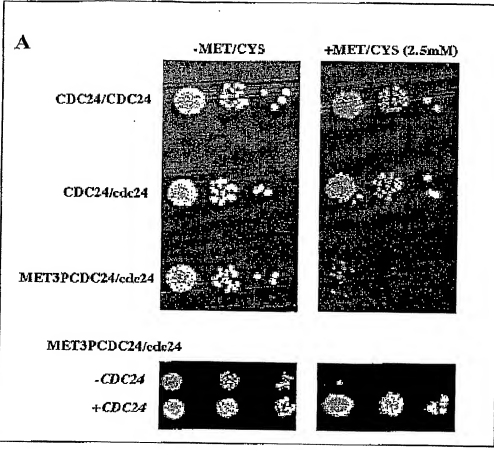
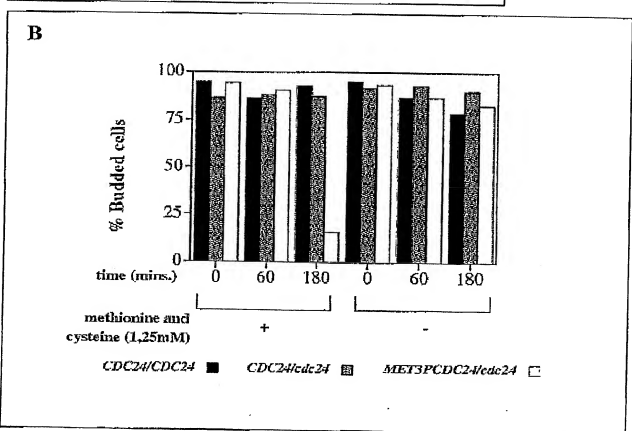


Figure 11











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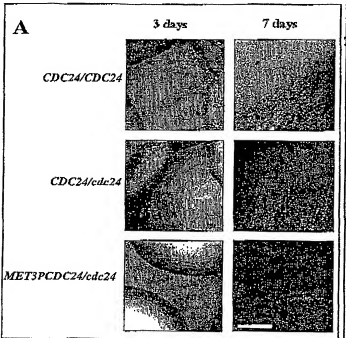
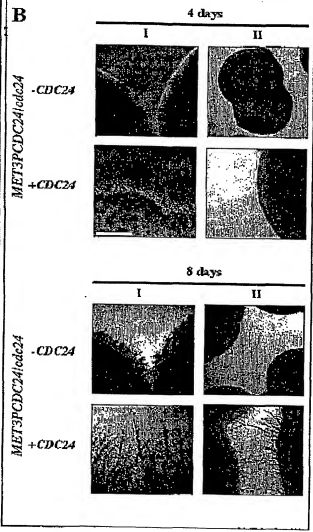
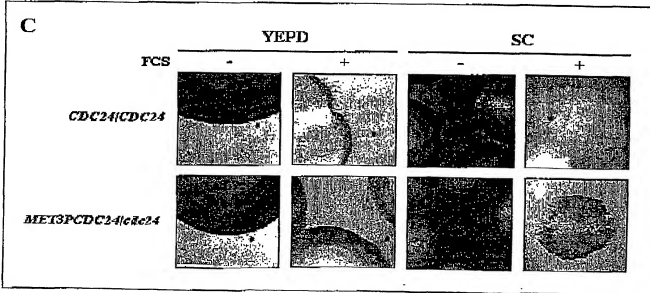
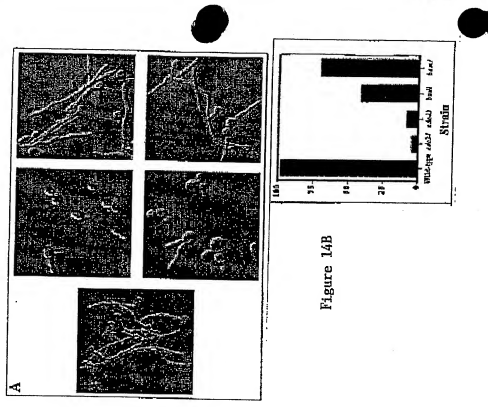


Figure 12







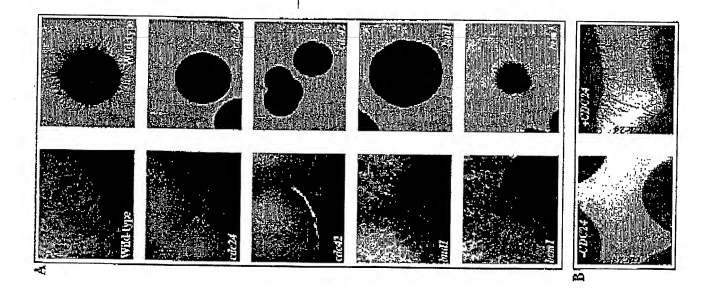


Figure 13

14A





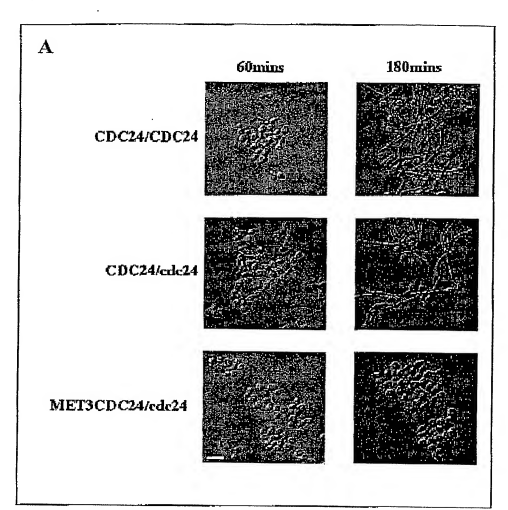


Figure 15

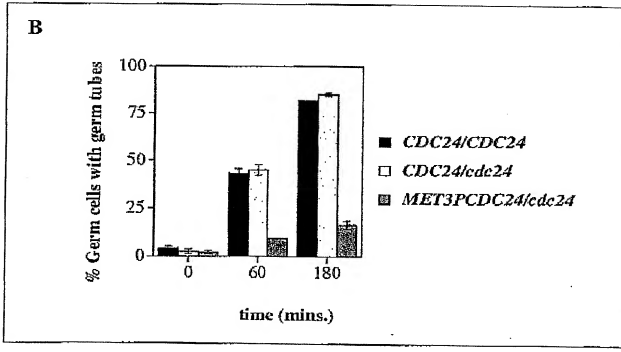




Figure 16

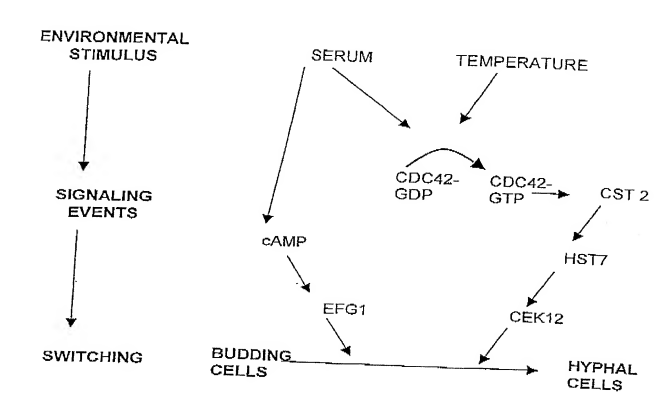




Figure 17

A. *cdc24* (wt)

5

SEQ. I.D. NO:1

DNA:

cccctctgtatacttttcaactctgtgaagccgcaatttaaattaccggtaatagcatctgacgatttgaaagtctgtaaaaaatccatttatgactt
tatattgggctgcaagaaacactttgcatttaacgatgaggagcttttcactatatccgacgtttttgccaactcgacgtcccagctggtcaaagt
gctagaagtagtagaaacgctaatgaattccagc

SEQ. I.D. NO:2

Protein:

15 PLCILFNSVKPQFKLPVIASDDLKVCKKSIYDFILGCKKHFAFNDEELFTISDVFANSTSQ LVKVLEVVETLMNSS

B. cdc24-m1

20 SEQ. I.D. NO:3

DNA:

cccctctgtatacttttcaactctgtgaagccgcaatttaaattaccggtaatagcatttgacgatttgaaagtctgtaaaaaatccatttatgacttt atattgggctgcaagaaacactttgcatttaacgatgaggagcttttcactatatccgacgtttttgccaactcgacgtcccagctggtcaaagt gctagaagtagtagaaacgctaatgaattccagc

25

SEQ. I.D. NO:4

Protein:

PLCILFNSVKPQFKLPVIAFDDLKVCKKSIYDFILGCKKHFAFNDEELFTISDVFANSTSQ LVKVLEVVETLMNSS



C. cdc24-m2

5 SEQ. I.D. NO:5

DNA:

cccctctgtatacttttcaactctgtgaagccgcaatttaaattaccggtaatagcatctggcgatttgaaagtctgtaaaaaatccatttatgactt tatattgggctgcaagaaacactttgcatttaacgatgaggagcttttcactatatccgacgtttttgccaactcgacgtcccagctggtcaaagt gctagaagtagtagaaacgctaatgaattccagc

10

SEQ. I.D. NO:6

Protein:

PLCILFNSVKPQFKLPVIASGDLKVCKKSIYDFILGCKKHFAFNDEELFTISDVFANSTSQ LVKVLEVVETLMNSS

15

D. cdc24-m3

SEQ. I.D. NO:7

DNA:

20 cccctctgtstacttttcaactctgtgaagccgcaatttaaattaccggtaatagcacctgacgatttgaaagtctgtaaaaaatccatttatgactt tatattgggctgcaagaaacactttgcatttaacgatgaggagcttttcactatatccgacgtttttgccaactcgacgtcccagctggtcaaagt gctagaagtagtagaaacgctaatgaattccagc

SEQ. I.D. NO:8

25 Protein:

 ${\tt PLCILFNSVKPQFKLPVIAPDDLKVCKKSIYDFILGCKKHFAFNDEELFTISDVFANSTSQ}\\ {\tt LVKVLEVVETLMNSS}$ 



SEQ ID NO. 10 STE4 DNA sequence (wild-type)

ATEGCACATCAGATGGACTCGATAACGTATTCTRATAATGTCACCCAACAGTATATACAACCACAAAGTCTACAGGA tacctctgcagcaggaagaattcaaaataaaatagaggcgccagacaagagagtaaacagcttcatgctcaaataa <u>ATRAAGCAAAACACAAGATACAAGATGCAAGCTTATTCCAGATGGCCAACAAAGTTACTTCGTTGACCAAAAATAAGATC</u> AACTTAAAGCCAAATATCGTGTTGAAAGGCCATAATAATAAAATCTCAGATTTTCGGTGGAGTCGAGATTCAAAACGTAT TTTGAGTGCAAGTCAAGATGGCTTTATGCTTATATGGGACAGTGCTTCAGGTTTAAAACAGAACGCTATTCCATTAGATT CTCAATGGGTTCTTTCCTGCGCTATTTCGCCATCGAGTACTTTGGTAGCAAGCGCAGGATTAAACAATAACTGTACCATT TATAGAGTTTCCAAAGAAAACAGAGTAGCGCAAAACGTTGCGTCAATTTTCAAAGGACATACTTGCTATATTTCTGACAT TGARTTTACAGRIRACGCRORTATATTGACAGCARGTGGGGRTRTGACATGTGCCTTGTGGGRTRTACCGRARGCARAGR GGGTGAGAGALTATTCTGACCATTTAGGTGATGTTTTGGCATTAGCTATTCCTGAAGAGCCAAACTTAGAAAATTCTTCG 10 ARCACATTCGCTAGCTGTGGATCAGACGGGTATACTTACATATGGGATAGCAGATCTCCGTCCTGTACAAAGCTTTTR ogranosni trong paratorno postorno properta de casa de caltara de caltara de caltara de caltara de como como c TRANTATGIA; CANTIAAGGICGGACTGIICTATIGCIACTTITICTCITTTTCGAGGTTATGHAGAACGIACCCCIRCC CCTACTTATATEGCAGCTAACATGGAGTACAATACCGCGCAATCGCCACAAACTTTAAAATCAATAAGCTCAAGCTATCT <del>reacaaccaagsogttetttttttagattttagt</del>gcatct<del>ggaagattga</del>tgfactcatgctatacagacatt<del>g</del>gtgte 15 ttgtgtgggatttattaaaggagagattgttggaaaattagaaggtcatggtggcagagtcatggtgtgcgctcgagi CCAGATGGGTTRGCTGTATGTACAGGTTCATGGGACTCAACCATGAAAATATGGTCTCCAGGTTATCAATAG

SEQ ID NO. 11

Ste4 Protein sequence (wild-type)

MAHQMDSITYSNNVTQQYIQPQSLQDISAVEEEIQNKIEAARQESKQLHAQINKAKHKIQDASLFQMANK/TSLTKNKII
LKPNIVLKGHNKKISDERWSRDSKRILSASQDGFMLIWDSASGLKQNATPLDSQWVLSCATSPSSTLVASAGLNNNCTI

25 RVSKENRVAQXVASIFKGHTCYISDIEFTDNAHILTASGDMTCALWDIPKAKRVREYSDHLGCVLALAIFEEPNLENSS:
TFASCGSDGYTYIWDERSPSAVQSFYVNDSDINALRFFKDGMSIVAGSDNGAIRMYDLRSDCSIATFSLERGYEEATPT.
TYMAANMEYNTAQSPQTLKSTSSSYLDNQGVVSLDFSASGRLMYSCYTDIGCVVWDVLKGEIVGKLEGHGGRVTGVRSS
EGLAVCTGSWDSTMKIWSPGYQ

30

Ý.

SEQ ID NO. 12 ste4-o15 DNA sequence (mutant)

RIGGCACATCAGATGGACTCGATAACGTATTCTAATAATGTCACCCAACAGTATATACAACCACAAAGTCTACAGGA

35 TATCTCTGCAGTGGAGGAGAAATTCAAAATAAAATAGAGGCCGCCAGACAAGAGGAGTAAACAGGTTCATGTTCAAAATT
ATAAAGCAAAACACAAGATACAAGATGCAAGCTTATTCCAGATGGCCAACAAGTTACTTCGGTGGAGTCGAGATTCAAAACGT;
AACTTAAAGCCAAATATCGTGTTGAAAGGCCATAATAATAAAATCTCAGATTTTCGGTGGAGTCGAGATTCAAAACGT;



SEQ ID No. 13

10

20

30

35

15 Ste4-o15 Protein sequence (mutant)

MAHQMDSITYSNNVTQQYIQPQSLQDISAVEEEIQNKIEAARQESKQLHAQINKAKHKIQDASIFQMANKVTSLTKNK
LKPNIVLKGHNNKISDFRWSRDSKRILSASQDGFMLIWDSASGLKQNAIPLDSQWVLSCAISPSSTLVASAGLNNNCT
RVSKENRVAQNVASIFKGHTCYISDIEFTDNAHILTASGDMTCALWDIPKAKRVRGYSDHLGDVLALAIPEEPNLENS.
TFASCGSDGYTYIWDSRSPSAVQSFYVNDSDINALRFFKDGMSIVAGSDNGAINMYDLRSDCSIAIFSLFRGYEERTP
TYMAANMEYNTAQSPQTLKSTSSSYLDNQGVVSLDFSASGRLMYSCYTDIGCVVWDVLKGEIVGKLEGHGGRVTGVRS:
DGLAVCTGSWDSTMKIWSPGYQ

SEQ ID NO. 14

25 ste4-o17 DNA sequence (mutant)



TAAATATGTATGATTTAAGGTCGGACTGTTCTATTGCTACTTTTTCTCTTTTTCGAGGTTATGAAGAACGTACCCCTACC
CCTACTTATATGGCAGCTAACATGCAGTACAATACCGCGCAATCGCCACAAACTTTAAAATCAACAAGCTCAAGCTATCI
AGACAACCAAGGCGCTGTTTCTTTAGATTTTAGTGCATCTGGAAGATTGATGTACTCATGCTATACAGACATTGGTTGTC
TTGTGTGGGGATGTATTAAAAGGAGAGATTGTTGGAAAATTAGAAGGTCATGGTGGCAGGTCACTGGTGCGCTCGAGI
CCAGATGGGTTAGCTGTATGTACAGGTTCATGGGACCCATGAAAATATGGTCTCCAGGTTATCAATAG

SEQ ID No. 15

Ste4-o17 Protein sequence (mutant)

- 10 MAHQMDSITYSNNVTQQYIQPQSLQDISAVEEEIQNKIEAARQESKQLHAQINKAKHKIQDASLFQMANKVTSLTKNKIN
  LKENIVLKGHNNKISDFRWSRDSKRILSASQDGFMLIWDSASGLKQNAIPLDSQWVLSCAISPSSTLVASAGLNNNCTI:
  RVSKENRVAQNVASIFKGHTCYISDIEFTDNAHILTASGDMTCALWDIPKAKRVREYSDHLGDVLALAIPEEPNLENSSI
  TFASCGSDGYTYIWDSRSPSAVQSFYVNDSDINALRFFKDGMSIVAGSDNGAINMYDLRSDCSIATESLFRGYEERTPT!
  TYMAANMEYNTAQSPQTLKSTSSSYLDNQGAVSLDFSASGRLMYSCYTDIGCVVWDVLKGEIVGKLEGHGGRVTGVRSS:
- 15 DGLAVCTGSWDSTMKIWSPGYQ
  - SEQ. I.D. No:16 refers to the epitope sequence "Tyr Pro Tyr Asp Val Pro Asp Tyr Ala".
  - SEQ. I.D. No:17 refers to TEV protease recognition sequence "Gln Asn Leu Tyr Phe Gln Gly"
  - SEQ. I.D. No:18 refers to peptide sequence "QFKLPVIAFDDLKVCKKSI".
  - SEQ. I.D. No:19 refers to peptide sequence "QFKLPVIASGDLKVCKKSI".
  - SEQ. I.D. No:20 refers to peptide sequence "QFKLPVIAPDDLKVCKKSI".
  - SEQ. I.D. No:21 refers to peptide sequence "QFKLPVIASDDLKVCKKSI".
  - SEQ. I.D. No:22 refers to peptide sequence "QYEFDVILSPELKVQMKTI".